

Identification of cell populations necessary for leaf-to-leaf electrical signaling in a wounded plant

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The identity of the cell files necessary for the leaf-to-leaf transmission of wound signals plants has been debated for decades. In *Arabidopsis*, wounding initiates the glutamate receptor-like (GLR)-dependent propagation of membrane depolarizations that lead to defense gene activation. Using a vein extraction procedure we found pools of GLR-fusion proteins in endomembranes in phloem sieve elements and/or in xylem contact cells. Strikingly, only double mutants that eliminated GLRs from both of these spatially separated cell types strongly attenuated leaf-to-leaf electrical signaling. *glr3.3* mutants were also compromised in their defense against herbivores. Since wounding is known to cause increases in cytosolic calcium, we monitored electrical signals and Ca^{2+} transients simultaneously. This revealed that wound-induced membrane depolarizations in the wild-type preceded cytosolic Ca^{2+} maxima. The axial and radial distributions of calcium fluxes were differentially affected in each *glr* mutant. Resolving a debate over which cell types are necessary for electrical signaling between leaves, we show that phloem sieve elements and xylem contact cells function together in this process.

jasmonate | Ricca's factor | xylem | phloem | calcium

Organ-to-organ electrical signaling is a highly conserved feature of land plants. For example, wound-induced electrical signals known as “slow wave potentials” (SWPs; otherwise known as “variation potentials”) have been found in numerous species (1, 2). In *Arabidopsis thaliana*, severe damage triggers electrical activity that propagates from leaf to leaf with apparent velocities in the centimeter-per-minute range. These events are characterized by rapid (<2 s) and massive (>50 mV) membrane depolarizations followed by slow (>5 min) repolarizations (3); each is a feature characteristic of SWPs (4, 5). At their destination in leaves distal to wounds, and strongly suggesting roles in plant defense, membrane depolarization in *Arabidopsis* stimulates the accumulation of the potent lipidic regulator jasmonate (JA). To initiate multiple defense responses, JA mediates the destruction of components of transcriptional repressor complexes to enhance gene expression (6–8). JA synthesis initiates this process. When a leaf is wounded, electrical signals travel to distal leaves to first trigger JA or JA precursor synthesis in a small population of cells known as xylem contact cells (9, 10). These cells then export JA precursors and/or JA to surrounding tissues (10). Which cells are necessary to conduct electrical signals to these distal sites of JA production?

Despite a debate that is decades old and that emerged in large part from studies of the sensitive plant *Mimosa pudica* (e.g., ref. 11), a consensus regarding which cell populations are necessary for organ-to-organ electrical signaling in plants has never been reached. On the one hand, the xylem has been proposed to play an essential, if not exclusive, role in this phenomenon. For example, a long-standing theory is that chemical elicitors (“Ricca's factor”) drawn over long distances through the xylem activate distal movement responses in *M. pudica* (12). This mechanism is thought to occur in other species (e.g., ref. 13). Alternatively, hydraulic signals in the xylem have been proposed to underlie SWP propagation (14). In each of these cases, dead xylem vessels rather than the living cells that surround them have been implicated in the

events leading to long-distance propagation of electrical signals. On other hand, evidence going back to the 1920s supports roles of the phloem in leaf-to-leaf electrical signaling (15–18), and wound-response electrical signals have been detected directly in sieve elements (19). Moreover, wounding initiates systemic calcium fluxes in *Arabidopsis* (e.g., ref. 20), and phloem cells have been implicated repeatedly in wound-response calcium signaling (17). Finally, both phloem and xylem-associated cells in *M. pudica* were found to be highly excitable (21), raising the possibility that both cell populations operate in the leaf movement response.

To address the question of which cell types are necessary for organ-to-organ electrical signaling in *Arabidopsis*, we based our approach on *GLUTAMATE RECEPTOR-LIKE* (GLR) genes (*GLR3.1*, *-3.2*, *-3.3*, and *-3.6*) that were recently found to function in this process (3). Functional GLR protein fusions were generated from three of these genes to identify the exact cell populations involved in SWP propagation. Then, based on the fact that wounding induces local and distal cytosolic calcium transients (20, 22), we developed a method to simultaneously monitor electrical activity and calcium levels. Our results build a cell-level understanding of the SWP.

Results

GLR Function in Membrane Repolarization and Defense Against Herbivores. The *glr3.3 glr3.6* double mutant attenuates wound-activated electrical signal propagation between leaves (3). To characterize additional genetic interactions and thereby define a

Significance

Numerous modes of long-distance electrical signaling exist in nature. The best known of these, axonal conduction, requires one primary cell population, i.e., neurons. In contrast, the cell types that mediate leaf-to-leaf electrical signaling in wounded plants have not been defined rigorously. Using genetic approaches, we find that two distinct populations of cells in the vasculature matrix are needed to perform this function. Surprisingly, these cells do not contact each other directly. As we further defined the plant wound response, we found that wound-induced membrane depolarizations preceded large intravascular calcium fluxes. We reveal a two-cell-type mode of electrical signaling in leaves and discuss parallels and differences in electrical signaling outside the plant kingdom.

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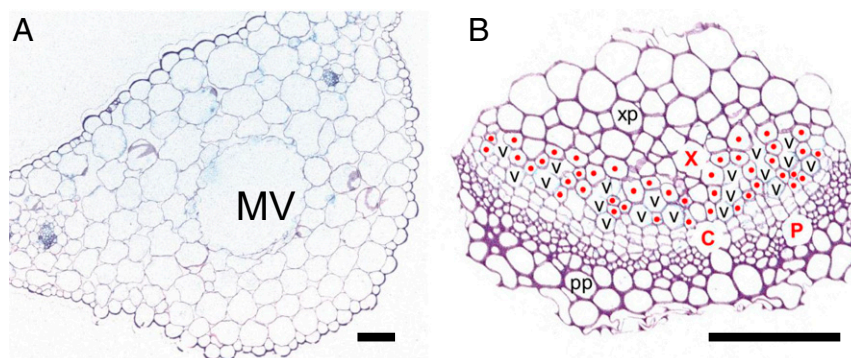


Fig. 3. Rapid midvein extraction from a 5-wk-old plant. (A) Transversal section near the leaf-petiole junction showing zone left by midvein (MV) after its rapid removal. (Scale bar: 30 μm .) (B) Transversal section of an extracted midvein. C, cambium region; P, phloem region; pp, phloem parenchyma; V, vessel; X, xylem region; xp, xylem parenchyma; red dots indicate contact cells. (Scale bar: 30 μm .) The sections were stained with 0.1% (wt/vol) toluidine blue in water.

Extracted midveins were either fixed and cleared directly (23, 24), subject to brief protoplasting to remove outer cell layers of parenchymal cells, or observed directly. Functional GLR3.1-VENUS fusion proteins localized primarily to XCCs (Fig. 4 A and B), and, in agreement with observation of the GLR3.1-GUS translational fusion (Fig. 2 G and F), a second low abundance pool of GLR3.1-VENUS was detected occasionally in phloem sieve elements (SEs) (SI Appendix, Fig. S1). In the root division zone, *GLR3.3* is expressed in a wide range of cell types (25). In the leaf primary vasculature, we found that GLR3.3-VENUS localized primarily to SEs (Fig. 4C). Sieve plate pores have been implicated as sites of ion channel expression (26). To observe sieve plates, isolated veins were subjected to complete protoplasting. In these samples, GLR3.3-VENUS was observed at the sieve plate periphery (SI Appendix, Fig. S2). Next, we examined GLR3.6-VENUS localization. Rapid vein extraction followed immediately by clearing and fixing revealed that GLR3.6-VENUS localized to xylem contact cells (SI Appendix, Fig. S3). When partial protoplasting was used to remove outer parenchymal cells of freshly extracted veins, the VENUS signal again located to large organelles within contact cells (Fig. 4D). In summary, phloem SEs and XCCs separated by the cambial region (SI Appendix, Fig. S4) contribute to propagation of the SWP from leaf to leaf.

Next, we assessed the major subcellular localizations of the GLR-VENUS fusion proteins. In the case of GLR3.1-VENUS, partial protoplasting to remove outer vascular cell layers was necessary to visualize the VENUS tag. GLR3.1 was found in punctate structures that resembled endoplasmic reticulum (ER). GLR3.1-VENUS was therefore introgressed into the red fluorescent protein (RFP)-ER marker line WAVE 6R (27). The fusion proteins and the ER marker proteins were found to colocalize in XCCs (SI Appendix, Fig. S5 A–C) and in the phloem SEs (SI Appendix, Fig. S5 D–F). In the case of GLR3.3-VENUS, punctate structures resembling ER were seen, and GLR3.3-VENUS was found to colocalize with the WAVE 6R ER marker (SI Appendix, Fig. S5 G–I). Crossing GLR3.6-VENUS into the WAVE 9R marker line that tags vacuolar membranes (27) revealed vacuolar colocalization of the two fluorescent tags (SI Appendix, Fig. S5 J–O).

Membrane Depolarization Maxima Precede Wound-Associated Cytosolic Calcium Maxima. GLR3.4 and GLR3.1/GLR3.5 heteromers form calcium-permeable channels (28, 29). Also, several GLRs control Ca^{2+} levels in pollen tubes (30), and *Physcomitrella* GLRs are Ca^{2+} -permeable channels (31). Furthermore, cytosolic Ca^{2+} levels increase in seconds in tissues proximal to wounds (22). With the goal of defining the temporal and spatial relationships of SWPs and Ca^{2+} signals we employed the GCaMP3 reporter (32)

that is known to detect Ca^{2+} fluxes in leaves (33). The presence of this protein expressed from the *UBIQUITIN 10* promoter did not strongly affect electrical activity after wounding (SI Appendix, Fig. S6). Additionally, each GCaMP3-expressing line was tested for its response to a nonwounding application of ice-cold water. The reporter responded similarly in each case (SI Appendix, Fig. S7). Experiments began with analysis of *Pieris brassicae* feeding on the WT background. For these initial analyses, WT plants at the 2-wk growth stage were chosen since calcium signals in veins are clearly demarcated by that age. As they fed, the larvae triggered vein-associated fluorescence (Fig. 5A and Movie S1). The apparent velocities of Ca^{2+} waves elicited in leaves distal to caterpillar feeding sites were greater in the petiole than in the laminal midvein and faster in the primary vein than in secondary or tertiary veins (Fig. 5B).

Since the three GLRs that we studied all localized to the primary vasculature, we tested whether the midvein alone was

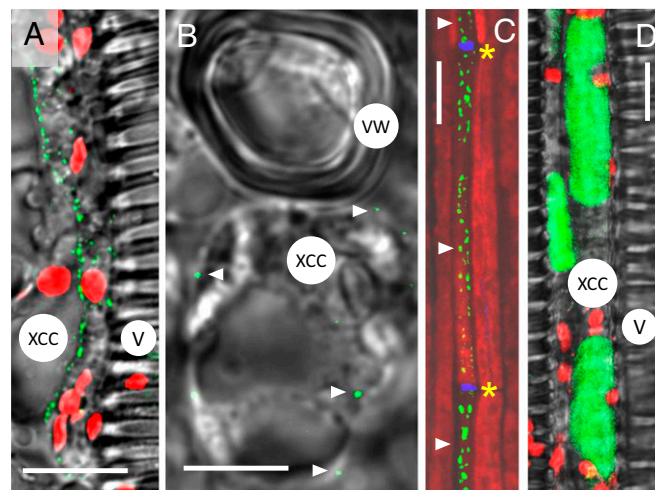


Fig. 4. GLR-VENUS expression in the primary leaf vein. (A) GLR3.1-VENUS localization (green) in contact cells; longitudinal section. Red is chlorophyll fluorescence. V, xylem vessel; XCC, xylem contact cell. (Scale bar: 10 μm .) (B) GLR3.1-VENUS (arrowheads) in a contact cell; transversal section. VW, vessel wall. (Scale bar: 5 μm .) (C) GLR3.3-VENUS localization (green) in SEs. The sample was fixed, cleared, and then stained with aniline blue and propidium iodide (red). Arrowheads indicate sieve cells; asterisks indicate aniline blue-stained sieve plates. (Scale bar: 10 μm .) (D) GLR3.6-VENUS (green) in xylem contact cells. Red is chlorophyll fluorescence. (Scale bar: 10 μm .) For A, B, and D mild protoplasting of freshly isolated veins in 500 mM sorbitol (A and B) or 100 mM sorbitol (D) was used to remove large peripheral parenchyma cells. For C, an isolated primary vein was fixed and cleared before staining.

the slower recovery phase correlated with the formation and diminution of the Ca^{2+} peak (Fig. 5D and Movie S2). Each *glr* mutant differentially reduced the amplitude of the Ca^{2+} wave in leaf 13. The *glr3.3* mutation, for example, attenuated peak GCaMP3 fluorescence in petioles more strongly than did the *glr3.1* or *glr3.6* mutations (Fig. 5E–G). Finally, in response to crush wounding, short duration surface potential changes were sometimes detected in *glr3.1 glr3.3*, but not in *glr3.3 glr3.6* (Fig. 5H and I). Fig. 5D–I and further analyses (Fig. 5J) showed that Ca^{2+} wave amplitudes correlated with electrical signal durations, and the apparent axial velocities of the two signals were similar.

Spatial analyses of GCaMP3 fluorescence after wounding revealed mutant-specific features (Fig. 5K–P). The Ca^{2+} signal in leaf 13 of *glr3.1* was often restricted to one leaf half-proximal to wounded leaf 8 (Fig. 5L). In *glr3.3*, the Ca^{2+} signal in this leaf was weaker and exclusively vascular compared with the signal from the WT, *glr3.1*, and *glr3.6* in which some perivascular GCaMP3 fluorescence was visible. Given the possible asymmetry in the Ca^{2+} signal in leaf 13 of *glr3.1* (Fig. 5L), we examined this mutant and found that the half of leaf 13 proximal to leaf 8 often produced stronger GCaMP3 fluorescence than did the leaf half distal to leaf 8 (SI Appendix, Fig. S11). This effect of the *glr3.1* mutation on the radial calcium signal is consistent with a different contribution to SWP propagation for GLR3.1 compared with GLR3.3 and GLR3.6.

Discussion

Fast nervous conduction in animals evolved under strong selection from predators (34). Here we argue that pressures from herbivores have led to the evolution of leaf-to-leaf electrical signaling in plants. It has long been known that mechanical stimulation of the sensitive plant *M. pudica* leads to membrane depolarization that results in visual crasis whereby leaves effectively disappear from view (15). Additionally, electrical signaling correlated with the production of defense-related proteinase inhibitors in tomato (35). More recently, genetic analysis revealed that electrical signaling stimulates the synthesis of the defense hormone JA in leaves distal to wounds. GLR3.3 is known to be necessary for full resistance of *Arabidopsis* to a fungus (36). Do the GLRs that participate in electrical signal propagation contribute to defense against chewing herbivores? This appears to be the case since the bioassays conducted in the present work (Fig. 1) show that larvae grown on plants carrying the *glr3.3* mutation gain weight more rapidly than they do on the WT. These findings underscore the importance of the GLR3.3 gene in defense activation and strongly implicate the phloem in signaling leading to activation of defense against herbivores. Together, our results support the hypothesis that a primary role of SWP in plants is to activate defenses in tissues distal to wounds. Defense-related electrical signaling is therefore common to both the plant and animal kingdoms.

Insights into Electrical Signaling. A key finding of our study was that, in the *Arabidopsis* SWP, two highly distinct vascular cell populations act to propagate wound signals from leaf to leaf. Without inactivating XCC-expressed GLR3.6 or GLR3.1 in the *glr3.1 glr3.3* and the *glr3.3 glr3.6* double mutants, *glr3.3* alone does not fully eliminate electrical signals in distal leaves. XCCs are perfectly placed to sense changes in solute composition or in water potential or water tension that may occur if xylem vessels are ruptured. Interestingly, natural variants of GLR3.6 in *Arabidopsis halleri* are found in environments that may affect plant hydration (37), and roles of GLRs in water pressure/tension sensing have been proposed (4). Strikingly, GLR3.6 shares a similar expression domain to lipoxygenase 6 (LOX6), which is necessary for the rapid synthesis of JA in leaves distal to wounds (9, 10). Therefore, contact cells play roles both in the propagation of JA-inducing electrical signals and in the synthesis of JA itself.

However, a unique role of xylem in leaf-to-leaf SWP transmission in *Arabidopsis* is untenable. This means that a xylem stream-transported Ricca's factor does not function as a unique leaf-to-leaf wound signal in our experiments. Both of the *glr* double mutants that attenuate electrical signaling harbor the *glr3.3* mutation. Confirming a crucial role of the phloem in leaf-to-leaf electrical signal propagation, GLR3.3 is expressed in SEs, which are among the most excitable cells in the plant (15, 21). The activation of the jasmonate pathway powerfully affects plant growth (38), and therefore JA synthesis must be controlled tightly. A two-tissue mechanism of signal propagation may assure tight control over defense activation in tissues distal to wounds. In summary, rather than using a single extravascular cell population for organ-to-organ electrical signaling, the plant SWP co-opts two distinct cell types in the vascular matrix. This mode of electrical signaling involves parallel, nonadjacent cell files.

Membrane Potential Changes Precede Ca^{2+} Maxima. Several clade 3 GLRs localize to organelle membranes and/or to the plasma membrane (28–30, 39–41). Our results do not rigorously identify pools of active GLRs, and we do not rule out the existence of functional GLRs in the plasma membrane. However, in our study, pools of GLR-VENUS fusion proteins in leaves localized preferentially to the endoplasmic reticulum or to the vacuole. This is of interest because a hyperactive variant of a calcium-permeable vacuolar cation channel powerfully activated JA responses (22). How this occurs is unknown, but local changes in cytosolic Ca^{2+} concentration and/or Ca^{2+} uptake from the cytosol were among the hypotheses proposed to link intracellular cation/ Ca^{2+} levels to JA synthesis activation (22). Moreover, SEs are sites of intense wound-response-associated calcium fluxes (17). Based on this, we simultaneously monitored cytosolic Ca^{2+} kinetics and electrical signals in leaves distal to wounds. By surgically removing extravascular tissues (Movie S2), we tested whether wound signals could be propagated through the primary vasculature. This was found to be the case for both Ca^{2+} signals and electrical signals. Therefore, signal propagation in leaves does not absolutely require extravascular cells. However, propagation of the calcium signal was delayed at the region of surgery. Either the integrity of the tissues surrounding the midvein helps to determine signal speed or epidermal GLR3.3 populations contribute to signal propagation.

A further finding was that each single *glr* mutant impacted the Ca^{2+} signal differently with *glr3.1* causing a bias in the radial distribution of the Ca^{2+} signal in the distal leaf. As with their impact on the electrical signal, the *glr3.1 glr3.3* and *glr3.3 glr3.6* double mutants strongly attenuated the distal calcium signal. We conclude that cytosolic calcium transients in response to wounding are regulated by several clade 3 GLRs and are strongly linked to the electrical signal. However, in the WT, GCaMP3 fluorescence maxima in leaves distal to wounds occurred on average 49 ± 8 s ($n = 7$) after the membrane depolarization maxima (e.g., Fig. 5D). How does this timing relate to JA accumulation in leaves distal to wounds? JA accumulation in leaves distal to wounds begins in the order of 90–120 s after wounding of distal leaves (9, 42). On average, it takes 66 s after wounding leaf 8 to initiate membrane depolarization in leaf 13 (3). Therefore, JA increases in the distal leaf lag behind the initial rapid depolarization phase of the SWP by at least 20 s. JA accumulation must therefore begin in or before the cytosolic Ca^{2+} maximum.

Comparison with Slow Wave Activities in Animals. Largely undefined at the cellular level, the plant SWP could not be readily compared with other electrical signaling phenomena in nature. However, the simultaneous monitoring of electrical signals and Ca^{2+} levels facilitates comparison of the SWP with electrical signaling phenomena found outside the plant kingdom. Such a comparison revealed several similarities to slow wave activities in

mammalian muscle such as that in the digestive system (43, 44): (i) rapid membrane depolarization and slow repolarization, (ii) centimeter-per-minute velocities of electrical signals, (iii) provocation of slow wave activity by current injection, and (iv) association with large and readily detectable cytosolic Ca^{2+} increases. While these parallels are intriguing, the mechanisms underlying plant and animal slow wave activities may differ since phloem and xylem are highly divergent from the cell populations involved in electrical signal propagation in metazoans. The deeper interest in clade 3 GLRs lies in the possibility that their study might yield novel insights into rapid communication between spatially separated cells.

Materials and Methods

All plants used were in the Col background. Generation of transgenics, complementation assays for GLR-VENUS fusions, insect bioassays, vein extraction, electrophysiology, and imaging procedures are documented in *SI Appendix, SI Materials and Methods*.

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